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- 64 Oily lubricant for cold plastic processing of metallic material.
- (f) An oily lubricant for cold plastic processing of metallic material obtained by compounding either one or both types of polymers of 2-ethyl polyacrylate-hexyl ester and polymethacrylic acid lauryl ester with one or more oiliness improvers selected from refined lard, oleic acid, phosphate, hindered ester, isostearic acid, and C₁₈ saturated higher alcohol and which are liquid at room temperature and which are compatible with the polymers. The oiliness improver is compounded as a thinning oil at a specified composition

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OILY LUBRICANT FOR COLD PLASTIC PROCESSING OF METALLIC MATERIAL

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The present invention relates to an oily lubricant for cold plastic processing of metallic material, and particularly to an oily lubricant used for cold plastic processing, such as drawing, contraction of area, rolling and forging of metallic material.

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For cold plastic processing, such as drawing, reduction in area (drawing), rolling, and forging, of metallic material, various forms of lubricants are added in order to prevent frictional damage (burning) of the metal mold, die, etc. and the material to be processed, such as steel pipe and steel sheet, and to improve the quality of the processed product as well as to reduce the wear of the tools.

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In conventional drawings oils and press oils acting as the oily lubricant, mineral oils or synthetic lubricating oils are used as the base oil, with oiliness improvers and extreme pressure additives added thereto. However, they are low in burning resistance; and when the degree of processing becomes high and the contact area between the material to be processed and the tool is increased, they tend to cause film breakage with resulting burning. Thus, they have only been useable for relatively light degree processing (for example, sinking), and it has been totally unfeasible to use them for steel pipe drawing, etc. under the severe processing conditions that exist. Also, since they contain extreme pressure additives, metal corrosion is caused.

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In conventional oily lubricants, as the synthetic lubricating oil acting as the base oil, polybutene, \angle -olefin,

oligomer, polyethyleneglycol, etc. are generally used, and as the oiliness improver, fats and oils, saturated and unsaturated fatty acids with 10 or more carbons, fatty acid esters, phosphate, alcohol, etc. are generally used; furthermore, as the extreme pressure additive, chemical compounds containing sulfur or chlorine are generally used.

The base oil is a carrier that has the function of dissolving the oily lubricant and the extreme pressure additive in a homogeneous and stable manner and to carry them into the portions needing to be lubricated. Mineral oils and sythetic lubricating oils do not have strong polar groups and they are low in lubricating ability. Accordingly, they cannot be used as the lubricant by themselves; however, they do not generally suffer chemical changes such as oxidation and deterioration, and thus, they are used as the base oil.

The oiliness improver is the material that is fed by the base oil to the metal friction surface requiring lubrication. and the lubricating effect is produced by the oiliness improver Fats and oils, fatty acids, etc. are long being adsorbed. chain compounds with high molecular weight, and they have a strong polar group at one end of the molecule. By the action of such polar group, the molecules are strongly adsorbed on and configured along the metal surface, thereby forming an Furthermore, because of the mutual adsorption film. attraction between the long molecular chains configured to make the adsorption film, the film becomes firm and strong, and exhibits a lubricating effect, to reduce the friction. However, as the temperature increases, the configuration of molecules become disordered, and at about 150°C, the film separates, and the lubricating effect is lost. oiliness improver loses its lubricating effect when the temperature reaches 150°C or above, the addition of an

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oiliness improver under severe lubricating conditions is not effective. Instead, an extreme pressure additive that exhibits the desired lubricating effect at high temperature is added.

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The extreme pressure additive is the material that is supplied, in the same manner as the oiliness improver, to the metal friction surface requiring lubrication, by the base oil, and forms a film on the metal surface, by chemically reacting with the metal when the degree of plastic processing becomes high and the contact area between the material to be processed and the tool becomes increased with the resulting extreme pressure condition which accompanies high temperature and high Through the formation of the film mentioned above, pressure. the extreme pressure additive prevents the burning by reducing the friction, and improves the burning resistance of the Chemical compounds containing chlorine or sulfur are different in reaction temperature range, respectively. Therefore, they are added in accordance with their purpose and use, when used together with the base oil and the oiliness For the extreme pressure additive containing chlorine, a chlorinated paraffin is used in many cases, and at 150°C to 250°C, the C-Cl bond is broken due to thermal decomposition, and Cl_2 or HCl is formed. Thus, the extreme pressure lubricating effect is shown. However, such extreme pressure additive has a disadvantage that, when water is present, hydrolysis is caused, and HCl is liberated, thereby causing serious corrosion. The boundary lubrication temperature range of the sulfur compounds is about 250°C to

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350°C.

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Also, as the conventional oily lubricants, there are those containing, in addition to the abovementioned three types of components, solid lubricants, such as graphite and molybdenum disulfide, which are mixed therewith in order to

improve the burning resistance. However, such oily lubricants tend to cause the burning, and in the same manner as the other oily lubricants, they can be used only for relatively low degree processing work (for example, sinking). Besides, such oily lubricants have disadvantages including the fact that the oil supply becomes difficult because of the presence of the solid lubricating agent, that they accumulate in the metal mold, or that they cause maintenance problems.

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As has been described above, in the use of the oily lubricant, by means of the base oil, the oiliness improver and the extreme pressure additive are supplied to the metal friction surface and adsorbed thereon. Then, in the temperature range from room temperature to about 150°C. as the fluid lubrication range in which the oiliness improver serves to maintain the oil film, the oiliness improver shows the desired lubricating effect; however, in the boundary lubrication range with higher temperature and with severe processing condition, the extreme pressure additive functions to maintain the oil film and exhibits a lubricating effect. However, in view of the low degree of the actual lubricating effect of the oily lubricant, due to its low burning resistance, the lubricant is only sufficient for low degree plastic processing for steel pipe drawing, etc.

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On the other hand, the oily lubricant can be used be merely applying it to the surface of the metal to be processed. Therefore, the process is simple and it is also quite easy to remove the lubricant after processing.

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Accordingly, various methods for improving the burning resistance of a lubricant as mentioned below have been employed conventionally.

For example, in the cold drawing of steel pipes, the following methods have been used:-

- a) The steel pipe is treated with a chemical conversion coating (for example, phosphate coating, oxalic acid oxidation coating) in advance, and is also provided with a secondary lubricant (for example, metallic soap, such as sodium stearate, and oil).
- 10 b) The steel pipe is coated with a liquid form synthetic resin (liquid obtained by emulsifying a synthetic resin, or by dissolving the synthetic resin in a solvent) in advance, and the coating is solidified by drying.
- Also, in the drawing of sheet form metallic material, the following methods have been used:-
- c) The lubrication film is thickened by directly coating the material with press oil that is made specifically high in viscosity.
 - d) Press oil with extreme pressure additive added thereto is used for directly coating the material.
- e) After coating the material with a solution of synthetic resin, such as vinyl chloride resin, hardening by drying is carried out.
- f) The synthetic resin film is directly adhered to the surface of the material to be processed.

However, these methods are not satisfactory in terms of simplicity in lubricating treatment, readiness for removal after processing, and the clearness of the metal surface.

From a practical point of view:

Method a) mentioned above enables the lubricant to show sufficient lubricating effect even when the degree of processing (draft) is high. However, since the lubricant is of the chemical reaction type, it is difficult to control the processing solution; and also because the life of the lubricant is short, it is necessary to carry out waste solution disposal treatment frequently. Thus, this method has an economic problem, and at the same time, it may cause environmental pollution. Besides, since the lubrication film adheres so well, it is difficult to remove the film from the product after processing. Furthermore, since the processing is of the chemical reaction type, and if the material is highly anticorrosive, the chemical treatment itself is difficult to carry out, and homogeneous chemical conversion coatings cannot be formed.

Method b) is to physically adhere the film over the surface of the metallic material, and a lubricant having lubricating characteristics superior to those obtained by method a) is produced. However, the method suffers from the disadvantage that it requires extensive drying in order to obtain the lubrication film. Also, the removal of the film after the processing becomes more difficult as the lubrication film becomes stronger.

Method c) is defective in that the processability becomes lower with an increase in viscosity; and

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Method d) is disadvantageous in that, since it uses a somewhat restrained corrosion reaction, severe corrosiveness is caused by the presence of water, etc.

Methods e) and f) make it possible to perform drawing of a high degree in draft and deep drawing which are not achievable by the drawing oil and press oil. However, they have the shortcomings in that they require strenuous work for drying and adhesion of the resin film, and also in that they are considerably higher in cost in comparison with the drawing oil and the press oil.

As has been mentioned above, while conventional lubricants have characteristic effects, respectively, every one of them has one or more disadvantages.

The present invention is intended to at least minimize the above-mentioned disadvantages of the prior art, and the object of this invention is to provide an oily lubricant for cold plastic processing of metallic material, that has a burning resistance nearly equal to that of lubricants obtained by conventional chemical conversion coating or synthetic film coatings, for the range from the fluid lubrication range to the boundary lubrication range.

Another object of this invention is to provide an oily lubricant for cold plastic processing of metallic material that does not cause corrosion to the metallic material.

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According to the present invention there is provided an oily lubricant for cold plastic processing of metallic material comprising one or both of the polymers selected from 2-ethyl polyacrylate-hexyl ester and polymethacrylic acid lauryl ester compounded with at least one type of an oiliness improver which is liquid at room temperature and which is compatible with said polymers, as a thinning oil, in a predetermined ratio.

We have found that an oily lubricant is obtained if the polymers of either one or both of 2-ethyl polyacrylate-hexyl ester and methacrylic acid lauryl ester are compounded with one type or not less than two types of oiliness improvers which are liquid at room temperature and also which are compatible with the foregoing polymers as a thinning oil, in a specified ratio. As suitable oiliness improvers which are liquid at room temperature and which are compatible with the polymers mentioned above, there may be mentioned refined lard, oleic acid, phosphate, hindered ester, isostearic acid, and \mathbf{C}_{18} saturated high alcohol, and mixtures thereof.

The inventors of this invention embarked on a study of an oily lubricant which is lowest in burning resistance among those applied in various types of lubrication processing methods, but which is able to simplify the process and is easily removable after the processing, in order to eliminate its corrosiveness to metals and to provide it with a burning resistance comparable to the oily lubricant obtained by conventional chemical conversion treatments or coating treatments, thereby making it feasible to actually apply it to the drawing of steel pipes.

First, the inventors decided to recheck the defective points of the conventional oily lubricants.

- (1) The corrosiveness to metals is derived from the extreme pressure additive contained in the oily lubricant; and as far as the extreme pressure additive is concerned, the corrosiveness to metals is unavoidable.
- (2) Mineral oils and synthetic lubricating oils used as the base oil are chemically stable at high temperatures and under high pressure, but they are extremely low in burning

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resistance. Because of the above, the oiliness improver and the extreme pressure additive are added.

Based on the above-mentioned understandings, the inventors started to discover, from high molecular weight compounds, a substance which is chemically stable (non-reactive), and which shows a far higher burning resistance (degree of oil film strength) than that shown by conventional ones, even in the boundary lubrication range (high temperature, high pressure), without using an extreme pressure additive.

The inventors expected that, of the high molecular weight compounds (polymers), those with markedly high molecular weight are high in chemical stability and suitable as the base oil, and also that some of them are high in burning resistance.

Accordingly, it was decided that, from innumerable high molecular weight compounds, those which do not need to be dissolved (those which are not in the form of a paste, powder, grain, or lump at room temperature) were to be selected. As a result, four types of substances which are liquid at room temperature were found, viz. (1) polybutene, (2) a copolymer of ethylene and \angle -olefin, (3) polybutadiene, and (4) butyl polyacrylic ester. Then by investigating their physical properties (average molecular weight and viscosity), the results shown in Table 1 were obtained. For these four types of substances there followed a Bauden test and a drawing test as the evaluation method for the lubrication characteristics, and the results obtained are shown in Tables 2 and 3. For the Bauden test, each sample agent was coated over the surface of the stainless steel (SUS 304). the test temperature was maintained at 100°C, a steel ball of 3/16' in diameter was pressed onto the coated surface

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with a 2 kg load, and it was made to reciprocate at a speed of 0.8 mm/sec. Under the foregoing conditions, the variation in the coefficient of friction of the sample agent was checked. Then, the coefficient of friction λ at the initial time and the number of frictions until reaching $\lambda = 0.2$ were evaluated as the burning resistance.

- 11 -Table 1 (Physical Property)

Sample Agent Tested	Average Molecular Weight (Mw)	Viscosity 100°C (cp)
1	2,350	4,000
2	3,600	2,000.
3	2,800	360
4	17,000	580

Table 2 (Bauden Test)

	Coefficient of Friction μ	Number of Times of Frictions until Reaching to $\mu=0.2$
1	0.198	.7
2	Higher than 0.2	0
3	0.168	10
4	0.160	Not less than 100

- 12 - Table 3 (Drawing Test)

÷.		1,	2	3	4
State of Pipe	Internal Surface	· x	0	. х	0
	External Surface	Δ	0	Δ	•
State of Plug		xx	0	x	0
Drawing Load (ton)	7.1	7.5	7.3	6.9
Plug Thrust (t	on)	1.6	1.4	1.4	1.2

Remarks:

- 1. Evaluation of the state of pipe
 - XX ... Burning is extensive.
 - X ... Slightly burnt.
 - Δ ... Metallic lustre and streak due to oil film breakage are observed.
 - O ... Slight streak.
 - 0 ... No abnormality is observed.
- State of plug
 - XX ... Burning is extensive.
 - X ... Slightly burnt.
 - O ... Cloudiness is observed.
 - O ... No abnormality is observed.
- Drawing load ... Indicated with maximum drawing load (gross drawing load).
- 4. Plug thrust ... Indicated with maximum plug thrust.
- 5. Tested under the drawing condition IV in Table 13.

In Table 2, the friction coefficient of (4) is 0.160, and this is considerably smaller than the general maximum threshold value 0.2 suitable for the lubricant. Also, the number of times of friction until the friction coefficient of exceeds 0.2 is remarkably large (100 or above). Also in Table 3, (4) shows a noticeably high value for the drawing test.

Therefore, it was decided that, of the tested agents (1) to (4), butyl polyacrylic ester of (4) alone was preferable.

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Thus, the inventors predicted that high molecular weight compounds having a molecular structure similar to that of butyl polyacrylic ester would be high in lubricating performance, and looked for ones which are in a liquid state at room temperature, selected from polyacrylic acids, polymethacrylic acids, and copolymers of acrylic acid esters and methacrylic acid esters.

Because polyacrylic acids are low in Tg (glass transistion temperature), although it depends on the number of carbons in the ester portion, many of them are in a liquid state at room temperature.

In general, polymethacrylic acids are high in Tg.

Accordingly, many of them are in powder form or solid state at room temperature. Of such methacylic acids, those found to be in a liquid state at room temperature were polymethacrylic acid lauryl ester and 2-ethyl polymethacrylate-hexyl ester.

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Since the copolymers of acrylic acid ester and methacrylic acid ester are either excessively viscous or in a solid state at room temperature, they were excluded by deciding that they are not suitable for the purpose of obtaining the liquid lubricant.

Table 4 shows the physical properties (average molecular weight and viscosity) of the selected high molecular compounds. Tables 5 and 6 show the results obtained for the drawing test conducted with regard to the burning resistance.

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As shown in Tables 5 and 6, every one of the high molecular compounds selected as shown in Table 4 was better than commercial products in results. Those showing particularly good results were (7) and (9).

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Table 4

No.	Sample Agent Tested	Average Molecular Weight (Mw)	Viscosity 100°C (cp
4	Butyl polyacrylate ester	17,000	580
5	Butyl polyacrylate ester	14,000	370
6	Ethyl polyacrylate ester	21,000	690
7	2-ethyl-polyacrylate -hexyl ester	14,000	470
8	2-ethyl-polymethacrylat	20,000	1,308,000
9	Polymethacrylic acid lauryl ester	48,000	4,420

Table 5

	Number of Sample Agent Tested		4	5	6	7	8	9	Commercial Product
-	****	*****	****	****	****	****	*****	****	****
Drawing Test	State of	Internal Surface	0	0	0	0	0	· Θ	0
	Pipe	External Surface	0	G	Θ	0	0	0	0
	State	of Plug	0	G	Ġ,	o	9	0	0
	Maximu Load (m Drawing (ton)	6.7	6.7	6.7	6.7	6.8	6.7	6.8
	Maximu Thrust	m Plug : (ton)	1.0	1.1	1.2	1.0	1.3	1.2	1.3

Remarks:

- Commercial product is G-755B metal working fluid (product name) from Nippon Kosakuyu Co., Ltd. A lubricating oil for drawing, that contains chlorinated paraffin as primary component; viscosity 100°C (cp) = 30.
- 2. The test was conducted under the drawing test condition I in Table 13.

Table 6

	umber of Sample gent · Tested		4	5	6	7	8	9	Commercial Product				
_	*****	******	****************										
Drawing Test	State of	Internal Surface	. 0	0	0	0	. 0	Θ	Δ				
- - - - -	Pipe	External Surface	0	0	0	0	9	0	0				
	State	of Plug	0	0	9	0	9	9	х				
	Maximu Load (m Drawing (ton)	7.0	7.0	7.1	7.1	7.2	7.0	7.3				
		m Plug (ton)	1.3	1.4	1.4	1.3	1.4	1.4	1.4				

Remarks:

- Commercial product is G-755B metal working fluid (product name) from Nippon Kosakuyu Co., Ltd. A lubricating oil for drawing, that contains chlorinated paraffin as primary component; viscosity 100°C (cp) = 30.
- 2. The test was conducted under the drawing test condition II in Table 13.

According to Tables 5 and 6, all of the high molecular compounds selected as listed in Table 4 are high in viscosity in comparison with the commercial products. Therefore, the inventors estimated that the single use of them would lower the workability. It is to be noted that in the ordinary operation, unless the kinematic viscosity at 50°C is 1,000 cst or below, the oil supply becomes difficult to use. Consequently, since 2-ethyl polymethacrylate-hexyl ester of (8) is extremely high in viscosity, it was thus evaluated to be impossible to put into actual use, and it was excluded.

Next, a study was conducted by aiming to lower the viscosity without lowering the lubricating performance (burning As the method to lower the viscosity, a thinning oil that is $150^{\,\mathrm{O}}\mathrm{C}$ or above in flash point and that is a low viscosity liquid at room temperature was considered. As the thinning oil, generally, an oiliness improver is used. As was mentioned in the beginning, as the oiliness improver, there are fats and oils, saturated or unsaturated fatty acid with ten or more carbons, fatty acid ester, phosphate, alcohol, etc., but those which are liquid with low viscosity at room temperature and which are 150°C or above in flash point are, for example, (a) refined lard, (b) oleic acid, (c) phosphate, (d) hindered ester, (e) isostearic acid, and (f) C_{18} saturated higher alcohol; and while (g) mineral oil is used as a base oil, it is generally used also as the thinning oil.

The thinning oils of (a) to (g) mentioned above are not usable practically as they are, and their compatability with the above-mentioned components (4), (5), (6), (7) and (9) which are to be compounded with them must be satisfactory. Therefore, all possible combinations between (4), (5), (6), (7) and (9) and (a) to (g) were taken into consideration,

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and they were compounded at a ratio of 1:1 in % by weight, and the compatability was checked. Thus, the results as shown in Table 7 were obtained.

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Then from Table 7, it was determined that (7) and (9) are high in compatibility with respective types of oiliness improvers (a) to (f) as well as with (g) and thus they are suitable for the purposes of this invention, and the range of the compounds to selected was narrowed down to 2-ethyl polyacrylate-hexyl ester of (7) and polymethacrylic acid lauryl ester of (9).

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The oiliness improvers are not limited to (a) to (f) mentioned above, but since they are liquids at room temperature and are compatible with the above mentioned polymers, they can be used as the oiliness improver in this invention.

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In addition, although the oiliness improver is compounded, its significance is different from that in conventional cases. As shown in Tables 5 and 6, since (7) and (9) show a high burning resistance by themselves, unlike in conventional synthetic lubrication oils, they are not to function as the base oil. The oiliness improver is added as a thinning oil.

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Table 7

37		·						
Number of Sample Agent Tested	4	5	6	7	8	9		
Mineral Oil (machine-10)	x	x	х	0		0		
Refined Lard	x	Δ	х	0		0	·	
Oleic Acid	0	0	х	0		0	•	
Phosphate	0	0	Δ	0		0		
Hindered Ester	Δ	Δ	Δ	0		0		
Isostearic Acid	Х	Δ	х	0		0		
C ₁₈ Saturated Higher Alcohol	х	X	Ō	0		0		

O ... Completely compatible
Δ ... Clouded liquid state, tended to separate
X ... Completed separated into two layers

Following the above, with each of 2-ethyl polyacrylate-hexyl ester of (7) and polymethacrylic acid lauryl ester of (9), (a) to (g) were compounded, respectively. By using them, many types of test oils were prepared by adjusting the compounding ratio so that the kinematic viscosity at 50°C becomes about 80, 150, 300, and 600 cst. Then, again, the compatibility and the burning resistance based on the drawing test were checked anew, and the results shown in Table 8, Table 9, Table 10 and Table 11 were obtained.

In Table 8, Table 9, Table 10 and Table 11, three types, that is (a), (b) and (c), are compounded with 2-ethyl polyacrylate-hexyl ester of (7) and polymethacrylic acid lauryl ester of (9), but it does not means to exclude (d) hindered ester, (e) isostearic acid, and (f) \mathcal{C}_{18} saturated high alcohol. Any oiliness improver that is liquid at room temperature and that is compatible with the above-mentioned polymers may be used.

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Table 8 (50°C Kinematic Viscosity about .80 cst)

		·				.80 CS	C)				- 1
NO. OF T	EST SAV	PLE OIL:		HI	H2 ·	н3	H4	RI	₹2	23	R4
% by wei	.gnt .cating	BOAI	. 7	43	. 29	38	46				
componer		S L	9			· .		29	16	20	30
		T O	h	.57	· · · · · · · · · · · · · · · · · · ·			71			
		IL OI N IM I L P	a		71				84.		
		N R G O	b			62				80	
	-	V E R	c				54				70
Compatil	oility			0	0	0	0	0	0	0	0
50°C Kin Viscosi		-		81.3	83.2	80.4	74.6	84.4	75.7	72.9	83.5
D T C	State of Pipe	Internal Surface		x	0	0	0	X	0	0	0
ASN		External Surface		Δ	. 0	0	0	Δ	0	0	0
I I	State	of Plug	·	х	O	0	0	٥.	0	0	0
G I 0 11	Maximu Load	m Drawing		7.0	6.9	6.9	6.85	7.1	6.9	6.8	6.85
II (ton	Maximu Thrust	m Plug		1.5	1.4	1.3	. 1.3	1.4	1.3	1.3	1.3
DTC REO	State of	Internal Surface		ХХ	0	. 0	0	XX	0	0	0
Y S Y C T W	2ipe	External Surface		Δ	0	0	0	۵	0	0	0
<u> </u>	State	of Plug		xx ·	۰ ٥	0	0	٥.	٥.	0	0
G I O N	Maximu Load	ım Drawing		7.4	7.2	7.0	6.99	7.5	7.2	7.2	. 7.3
III (tor	Maximu Thrust	ım Plug		1.7	1.5	1.6	1.5	1.6	5 1.5	1.6	1.55
											

Remarks: 1. The compatibility is indicated by the same manner as in Table 7.

2. The test was conducted under the Drawing Test Conditions II and III of Table 13.

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<u>Table 9</u> (50°C Kinematic Viscosity about 80cst)

 ====		(2) - 2 -										
NO. OF T	EST SAM		•		ਜ5	н6 -	<u>H7</u>	H8	_R5	R6	R7	R8
<pre>\$ by wei of lubri</pre>	cating	A I		. 7	53	43	51	57				
componen	ts	S L E		9					35	25	29	38
		TO		'n	46	•			65			
	•	IL (I M L P	a		57 [°]	•		•	75		
	-	N G	я 0	ď			49				71	. •
		·	V E R	С				43				62
Compatibility					0	0	0	0	0	0	0	٥
50°C Ki: Viscosit					147	149	155 _. .	152	150	151	149	154
D T C	State of . Pipe	Intern Surfac	2		Δ	0	• Θ	e	Δ		9	G
A S M W T D		Extern Surfac			0	•	9	. 0	0	9	G	0
I I	State	State of Plug Maximum Drawing. Load			x	0	Θ.	9	х.	•	9	Θ
й С <u>т</u> Ой	Maximu Load				7.0	6.8	6.9	6.8	7.1	6.9	6.9	6.8
II (ton	Maxinu Thrust	m Plug			1.45	1.3	1.4	1.4	1.4	1.3	1.3	1.3
D T C	State	Intern	:2		x	0	0	0	x	0	0	0
	?ipe	Exter: Surfac			0	9	9	9	0	9	Θ	G
A S D I E I I O	State	of Plug	ī		х	0	G	Ø -	х.	0	9	G
G I	Maximu Load	Maximum Drawing Load			7.4	7.2	7.4	7.3	7.5	7.3	7.2	7.2
III (ton) Maximum Plug Thrust					1.7	1.55	1.6	1,55	1.75	1.6	1.6	1.5

REMARKS: 1. The compatibility is indicated in the same way as in Table 7.

2. The test was performed by using the drawing test conditions II and III in Table 13.

- 23 - Table 10 (Drawing Test)

									,	,			
NC. OF	TEST SA		OIL:		Н9	H10	HIL	н12	R9	R10	R11	R12	
	icating	A I	_	. 7	65	57	62	68					С 0
comboue	nts	S L E		9	-				43	35	38	46	N P
		H I		h	35				57				V R E O
-		N I	I M L P	a	•	43	•	:		65			N D T U I C
		ห G	R 0	b			38				62	- ,	O T
			V E R	c				46				- 4	A L
Compati	bility				0	0	0	0	ο .	0	0	0	
50°C Ki Viscosi				-	326	283	298	313	318	292	306	302	290
D T C R E O A S N	State of Pipe	Inte	rnal ace		Δ	0	0	0	Δ.	9	G	G .	0
A S N W T D		Exte Suri			0	0	9	Θ	. 0	0	.0	G	_Q
N T D	State	of 71	ug		0	0	9	0 .	0	Θ	• 0	0	0
G I O N	Maximu Load	m Dra	wing		7.0	6.8.	6.8	6.8	7.1	6.8	6.9	6.8	7.0
II (ton) _{Maximu} Thrust	m Plu		•	1.65	1.5	1.5	1.6	1.5	1.45	1.5	1.4	1.5
DTC REO	State	Surf		•	XX	· 0	0	0	хх	0	0	0	•
ASX W T D	Pipe	Exte	rnal ace		Δ	0	0	o,	Δ.	0	0	0	0
7 7 7	State	o : 71	.ug		хх	0	0	0	0.	0	0	0	х
3 9 1	Maximu Load	um Dra	wing		7.4	7.2	7.0	6.95	7.5	7.2	7.2	7.3	7.4
III (tor	Maximu Thrust	ım Plu	ıg		1.7	1.5	1.6	1.5	1,65	1.5	1.6	1.55	1.65

Remarks: 1. The compatibility is indicated in the same way as in Table 7.

- The test was performed by using the drawing test conditions II and III in Table 13.
- 3. The conventional product used for comparison is G-755B (product name) from Nippon Kosakuyu Co., Ltd. It is a lubricating oil for drawing, containing chlorinated paraffin as the main component. 100°C 30 cp.

- 24 <u>Table 11</u> (Drawing Test)

		PLE OIL:		H13	H14	H1:5	H16	R13	Rl4	R15	R16
% by wei		3 O A I	. 7	78	70	77	80				
componen			9			 		50	44	48	53
		T O	h	22			-	50			
		IL OI N IM I LP N R	a		30	•	•		56		
			þ			23		-		52	
		V E R	c	÷	•		2,0				47
Compatib	ility	-		0	0	0	0 .	o	0	0	0.
	50°C Kinematic. Viscosity cst				614	685	611	565	592	605	634
DTC	State of	Internal Surface		Δ	· O	Θ.	0	Δ	G	0	9
ASM WID	Pipe	External Surface		0	0	9	9	0	Θ	Ð	Э
7	State (of Plug		0	0	. 0	9	٥ .	Θ	Э	9
G I O	Maximu Load	m Drawing		7.1	6.9	6.8	. 6.8	7.25	6.8	6.8	6.8
II (ton	Maximu Thrust	m Plug		1.45	1.4	1.45	1.4	1.55	1.45	1.45	1.4
3 E O 3 E O 4 S N	State	Internal Surface		х	9	0	0	x	Э	Θ	0
כיי א	Pipe	External Surface		0	9	9	0	0	0	0	0
I I N T G I	State	of Plug		xx	Э	Θ	0	ХХ	. 0	G	<u> </u>
	Maximu Load	num Drawing		7.5	7.3	7.3	7.3	7.55	7.2	7.3	7.25
III (ton	III (ton) Maximum Plug Thrust				1.55	1.6	1.5	1.75	1.6	1.6	1.5

Remarks: 1. The compatibility is indicated in the same way as in Table 7.

The test was performed by using the drawing test conditions II and III in Table 13. It was found from Tables 8 to 11 that burning is caused less on the external and internal surfaces of the tested materials (drawn pipes) when the oiliness improver is compounded as the thinning oil, than that caused in the lubrication performance shown by the case where the respective polymers are used as the only component. In particular, those diluted with oleic acid or phosphate are satisfactory in lubricating performance, and also, those with higher viscosity are proportionally higher in performance. In the case of 80 cst, slight burning was observed. The case in which mineral oil was compounded as the thinning oil was unsatisfactory in performance.

Next, the burning resistance was checked for the samples obtained by diluting them with a thinning oil (equal % by weight compounding) prepared by combining not only one type but not less than two types selected from lard, oleic acid and phosphate.

The thining oils (equal % by weight compounding) were prepared by combining not less than two types selected from lard, oleic acid and phosphate, and the respective thinning oils thus obtained were compounded (also in this case, the compounding ratio was equal to each other among the components in terms of % by weight) with 2-ethyl polyacrylate-hexyl ester or polymethacrylic acid lauryl ester. Also, an adjustment was made for the kinematic viscosity so that it became about 300 cst at 50⁰C. In this manner, many types of test oils were prepared. Together with the products (lubricants) obtained by using the conventional art, the test oils were tested by the drawing test under the drawing test condition IV in Table 13, and the results shown in Table 12 were obtained. The drawing test condition IV is 45.9% in reduction of area, and this is a processing degree (draft) close to the limit for the drawing of carbon steel pipe, for

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one time. If the draft is increased even slightly from it, the material becomes ruptured.

From the results shown in Table 12, it was found that the lubricants provided by this invention have a burning resistance almost equivalent to that shown by the conventional lubricants (those obtained by the chemical conversion coatings and by the synthetic coating processing). Also, it was found that when the metal face of the outer surface of the steel pipe after the drawing conducted by using the lubricants according to this invention was compared with such metal surface drawn by using the conventional lubricant, the metal surface resulting from the use of the lubricant according to this invention is cleaner than that resulting from the use of the conventional lubricant.

From the results obtained as mentioned above, it was confirmed that the samples prepared by compounding the thinning oil that is composed of a single component selected from oleic acid, lard and phosphate which are the oiliness improvers, or that is composed of two or more types of components selected from the foregoing oiliness improvers, with either 2-ethyl polyacrylate-hexyl ester or polymethacrylic acid lauryl ester have excellent burning resistance comparable to that shown by lubricants obtained by chemical conversion coatings or resin coatings.

Furthermore, since 2-ethyl polyacrylate-hexyl ester and polymethacrylic acid lauryl ester are compatible with each other, the thinning oil composed of a single component or the combined components selected from oleic acid, lard and phosphate may be compounded with the compound of 2-ethyl polyacrylate-hexyl ester and polymethacrylic acid lauryl ester.

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	50° Kinema	on) Viscosity est	310	301	301.	Ċ	887	297	292	308	1	285	1		1	
	ΙΞ	Load (ton)	2.4	2.2	2.35	L	2.45	2.45	2.2	2.3		. 2 . 4	2.1		2.2	
Test)	Maximum Stretching	Load (ton)	7.7	7.6	7.8	1	7.8	7.75	7.6	7.65		7.8	7.6		7.65	
(Drawing	State of	Plug	0	G	O .	•	0	0	0	0		o .	0	٠	0	
Table 12 (Pipe External	Surface	0	0	0		©	0	0	•		•	③		•	
Ä	 44 -4	Surface	0	•	0	,	0	0	0	0		0	0		•	
	Base	Oil	7	7	7	ı	7	6	6	6		O				
	Composition of Thinning	0i1	Oleic Acid & Lard	Oleic Acid & Phosphate	Lard & Phosphate	Oleic Acid	Lard & . Phosphate	Oleic Acid & Lard	Oleic Acid & Phosphate	Lard & Phosphate	Oleic Acid,	Lard, & Phosphate	Phosphate as base & Sodium	Stearate	Resin Coating	rocessing
	No. of Oil	Tested	H17	H18	H19		H20	R17	R18	R19	-	R20	PRIOR	ART	•	

Remarks:

The compatibility is indicated in the same way as in Table 7.

^{2.} Base oils (7) and (9) are those in Table 4.

Thinning oils ((h), (a), (b), (c), (d) are those in Table 8. m m

Table 13 (List of Drawing Test Conditions)

		Sample Structure 0.2% C Matter Master Pipe Dimen Tested wall thickness 2. Oil Supply: Imme	(STKM-13C) sions: Out 5mmt		
5	Condition	Dimensions after drawing (outside diameter x wall thickness)	Reduction in Area -	Plug	Drawing Rate
	I	19.00mmø x 2.3mmt	32.9%	Spherical Plug	10m/min
10		19.00mmø x 2.1mmt	38.0% -	Spherical Plug	10m/min
	III .	19.00mmø x 1.9mmt	43.2%	Spherical Plug	10m/min
15	IV ·	19.00mmø x 1.8mmt	45.9%	Spherical Plug	10m/min

As seen in Tables 8 to 11, when the mineral oil is added, the lubricating performance of the polymers is lowered. However, for example, in the area reduction processing for metal sheet, compared with the drawing of steel pipe, the lubricating condition is considerably Therefore, if an extremely low cost mineral oil can be added without causing a significant lowering of the lubricating performance, it is possible to use the mineral oil as a filler (extending agent), and thus, it is economical. Accordingly, with respect to the lubricating performance of the polymers in terms of the burning resistance in cold plastic processing, the possibility of mixing mineral oil to make it feasible to carry out cold plastic processing with a burning resistance that is higher than that shown by conventional oily lubricant although it is lower than that shown by the conventional chemical conversion coating lubrication and synthetic resin coating lubrication was investigated.

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For the purpose mentioned above, the inventors prepared test oils by further compounding mineral oil, at various ratios, with the oily lubricant obtained by compounding the above mentioned thinning oil with l-ethyl polyacrylate-hexyl ester. Then, by conducting an area reduction (drawing) test, the relation between the lowering in lubricating performance and the ratio of the mineral oil compounded was checked. The results are shown in Table 14. As is seen in this Table, each test oil has a lubricating performance range best suitable for the area reduction (drawing) processing of metal sheet, that is determined in accordance with the compounding ratio.

Table 14 (Area Reduction Test)

	Blank Diameter mm Ø	eter mm ø		92	93	94	95	96		86	66	100
	Reduction Ratio = blank diameter + punch diameter	Ratio ameter ameter	-	2.3	2.325	2.35	2.375	2.4	2.425	2.45	2.475	2.5
%q %	Oily Lubricant	Mineral Oil							•			
weight	100 parts	0 parts	149					0'0	0,0	0,0	x,x	x,x
o f	100 parts	10 parts	9 5				0,0	0,0	о'х	х,х		
test	100 parts	20 parts					0,0	0,0	х'х			-
oil	100 parts	40 parts	001			0,0	0,0	×,×	•	•	-	•
	100 parts	60 parts	CSC		0'0	0,0	x,x					
CONVEN- TIONAL	Commercial Press Oil		160			0'0	0'0	x, x	x,x	•		
PRODUCT	Resin Coating Lubrication	ing n						0,0	0,0	0'0	х о	x,x
		50 v1	50°C kinematic viscosity cst	nemati ty cst	v							•

Table 14 (Area Reduction Test)

- A. Remarks:
- Cylinder drawing (area reduction) test condition
 Punch diameter; 40 mm Ø, shoulder 4.5 mm R
- Die diameter; 42.58 mm ø, shoulder 9.1 mm R

 Sample material tested; spec 0.8 mmt (cold rolled steel sheet)

 Drawing rate; 20 M/min

 Wrinkle press; 700 kg
 - 2. O ... Drawing is succeeded, X ... Ruptured
- 10 3. cst indicates the kinematic viscosity at 50°C.
 - 4. For the product according to this invention, the sample H6 in Table 9 was used.
 - 5. As the thinning oil, the machine oil 220 that is similar to H6 in viscosity was used.
- 15 6. As the commercial press oil, No. 640 press machine oil (metal working fluid) produced by Nippon Kosakuyu (50°C 160 cst, with chlorine, sulfur system extreme pressure additive compounded) was used.
- 7. For the resin coating lubrication processing, a resin coating type lubricant, commercial name "Lubcoat N-7" from Nippon Kosakuyu, was used.

Finally, for the sample materials for the drawing and area reduction tests using the test oils according to this invention obtained by varying the compounding ratio of the components mentioned above, respectively, the following test was conducted. That is, the sample materials were immersed in an aqueous solution of 3 % sodium orthosilicate that was heated to 60° C. The result is that every sample could be freed from grease completely. Also, no discoloration or corrosion was caused to the steel material.

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Ash should be apparent from the description given above, the oily lubricant for cold plastic processing of metallic material, that is provided by this invention, has the following effects.

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(1) While it is an oily lubricant, it has a burning resistance equivalent to that shown by the chemical conversion coating lubrication and the synthetic resin coating lubrication. Consequently, various types of cold plastic processings in which the quality of the resulting product was assured by the chemical conversion coating lubrication and the synthetic resin coating lubrication can employ the direct oil supply system instead of the above mentioned coating lubricants. Thus, the lubricating treatment becomes simple and less troublesome. As a result, the process becomes simpler than the process using the chemical conversion coating lubrication and the resin coating lubrication, and such direct oil supply system can be applied also to the continuous processes. Furthermore, various types of cold plastic processing products which have been assured of their quality by chemical conversion coating lubrication and synthetic resin coating lubrication are easy to treat for the removal of lubricating oil after processing due to the use of the oily lubricant.

metal surface becomes cleaner than that of products processed by using chemical conversion coating lubrication or synthetic resin coating lubrication. Also, when lubricating performance can be a little lower, such as in the area of reduction processing for metal sheet, mineral oil can be used as a filler.

(2) As it contains not only extreme pressure additives, but also other materials which are high in reactivity, the metal material does not corrode, thus, the defects of oily lubricants provided by prior art can be eliminated.

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CLAIMS

1. An oily lubricant for cold plastic processing of metallic material comprising one or both of the polymers selected from 2-ethyl polyacrylate-hexyl ester and polymethacrylic acid lauryl ester compounded with at least one type of an oiliness improver which is liquid at room temperature and which is compatible with said polymers, as a thinning oil, in a predetermined ratio.

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2. An oily lubricant for cold plastic processing of metallic material as set forth in claim 1, wherein said oiliness improver is selected from lard, oleic acid, phosphate, hindered ester, isostearic acid, or \mathbf{C}_{18} saturated higher alcohol and mixtures thereof.

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